chain nodes:
6 7 8 9 10 11 12 14 16 17 18
ring nodes:
1 2 3 4 5
chain bonds:
1-16 2-18 4-6 6-7 6-11 7-8 7-17 8-9 8-12 9-10 9-14
ring bonds:
1-2 1-5 2-3 3-4 4-5
exact/norm bonds:
1-2 1-5 2-3 2-18 3-4 4-5 6-7 6-11 7-8 8-12 9-10 9-14
exact bonds:
1-2 1-5 4-3 7-18 8-9

G1:H, CH3, CH2, CH, Et, n-Pr, i-Pr, n-Bu, i-Bu

G2:0,N

Match level: 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:CLASS 7:CLASS 8:CLASS 9:CLASS 10:CLASS 11:CLASS 12:CLASS 14:CLASS 16:CLASS 17:CLASS 18:Atom

L1 STRUCTURE UPLOADED

=> d L1 HAS NO ANSWERS L1 STR

G1 H, Me, CH2, CH, Et, n-Pr, i-Pr, n-Bu, i-Bu G2 O, N Structure attributes must be viewed using STN Express query preparation.

=> s 11 full

FULL SEARCH INITIATED 14:09:01 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 2491 TO ITERATE

100.0% PROCESSED 2491 ITERATIONS 24 ANSWERS SEARCH TIME: 00.00.01

24 SEA SSS FUL L1

=> file caplus

COST IN U.S. DOLLARS SINCE FILE TOTAL. ENTRY SESSION FULL ESTIMATED COST 178.36 178.78

FILE 'CAPLUS' ENTERED AT 14:09:09 ON 11 FEB 2008 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.

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http://www.cas.org/infopolicy.html

=> s 12 T.3

6 L2

=> d 13 1-6 ibib abs hitstr

L3 ANSWER 1 OF 6 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 2006:817369 CAPLUS Full-text

inhibitors

DOCUMENT NUMBER: 145:249516

TITLE: Preparation of peptide boronic acids as proteasome

INVENTOR(S): Oliva, Ambrogio; Bernardnini, Raffaella; D'Arasmo, Germano; Cassara, Paolo G.; Bernareggi, Alberto;

Menta, Ernesto

PATENT ASSIGNEE(S): Cephalon, Inc., USA SOURCE: PCT Int. Appl., 159pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	PATENT NO.					KIND		DATE		APPLICATION NO.								
WO	WO 2006086600			A1		20060817		WO 2006-US4664										
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		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	KP,	KR,	
		KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	
		MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	
		SG,	SK,	SL,	SM,	SY,	TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	
		VN,	YU,	ZA,	ZM,	ZW												
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		GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,	
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CA	2597	273			A1		2006	0817		CA 2	2006-	2597	273		2	0060	210	
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										US :	2006-	3511	93		A 2	0060	209	
										wo :	2006-	US46	64		W 2	0060	210	
OTHER S	HER SOURCE(S):					MARPAT 145:2495												

GI

AB The invention provides peptide boronic acid derivs. Hy-CONHCHR2CONHCH(CH2CHMe2)B(OR1)2 [R1 is H, alkyl, cycloalkyl, cycloalkylalkyl, aryl, aralkyl or may combine to form a ring; R2 is CHMeOH or aminomethyl; Hy is an optionally-substituted nitrogen-containing heterocyclic group optionally fused with an aryl or heteroaryl group (with provisos) | that can modulate apoptosis by inhibition of proteasome activity and are for use in treating diseases such as cancer and other disorders associated directly or indirectly with proteasome activity. Thus, compound I was prepared by a multistep sequence starting with reaction of (+)-pinanediol with 2-methylpropylboronic

acid, conversion of the product to a leucine boronate analog, and subsequent acylations by Boc-protected L-threonine and 6-phenyl-2-pyrazinecarboxylic acid.

IT 906089-74-5P 906090-30-0P 906090-65-1P 906090-90-2P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of peptide boronic acids as proteasome inhibitors)

RN 906089-74-5 CAPLUS

CN 1H-Pyrazole-3-carboxamide, N-[(1S,2R)-1-[[(1R)-1-[(3aS,4S,6S,7aR)-hexahydro-3a,5,5-trimethyl-4-6-methano-1,3,2-benzodioxaborol-2-yl]-3-methylbutyl]amino]carbonyl]-2-hydroxyropoyl]-5-phenyl- (CA INDEX NAME)

Absolute stereochemistry.

- RN 906090-30-0 CAPLUS
- CN 1H-Pyrazole-3-carboxamide, N-[(1S)-2-[[(1R)-1-[(3aS,4S,6S,7aR)-hexahydro-3a,5,5-trimethyl-4,6-methano-1,3,2-benzodioxaborol-2-yl]-3-methylbutyl]amino]-1-[[(4-methylbenzoyl)amino]methyl]-2-oxoethyl]-5-phenyl-(CA INDEX NAME)

Absolute stereochemistry.

- RN 906090-65-1 CAPLUS
- CN Boronic acid, [(1R)-1-[[(2S,3R)-3-hydroxy-1-oxo-2-[[(5-phenyl-1H-pyrazol-3-y1)carbonyl]amino]butyl]amino]-3-methylbutyl]- (9CI) (CA INDEX NAME)

RN 906090-90-2 CAPLUS

CN Boronic acid, [(1R)-3-methyl-1-[[(2S)-3-[(4-methylbenzoyl)amino]-1-oxo-2-[[(5-phenyl-1H-pyrazol-3-yl)carbonyl]amino]propyl]amino]butyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 2 OF 6 CAPLUS COPYRIGHT 2008 ACS on STN 2005:216605 CAPLUS Full-text

ACCESSION NUMBER:

DOCUMENT NUMBER: 142:316496

TITLE: Preparation of substituted cycloalkylamine derivatives as modulators of chemokine receptor activity

INVENTOR(S): Carter, Percy H.; Cherney, Robert J.; Batt, Douglas G.; Brown, Gregory D.; Duncia, John V.; Gardner,

Daniel S.; Yang, Michael G.

PATENT ASSIGNEE(S): Bristol-Myers Squibb Company, USA SOURCE:

PCT Int. Appl., 440 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA	TENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D	ATE	
		50208 50208			A2 A3		2005			WO 2	004-	US27	195		2	0040	320
	W:	GE, LK,	CO, GH, LR,	CR, GM, LS,	CU, HR, LT,	CZ, HU, LU,	AU, DE, ID, LV, PL,	DK, IL, MA,	DM, IN, MD,	DZ, IS, MG,	EC, JP, MK,	EE, KE, MN,	EG, KG, MW,	ES, KP, MX,	FI, KR, MZ,	GB, KZ, NA,	GD, LC, NI,

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            SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,
            SN, TD, TG
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    US 2005054626
                         A1
                               20050310
                                                                  20040819
    EP 1656138
                         Α2
                               20060517
                                           EP 2004-781805
                                                                  20040820
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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                               20070215
    JP 2007502842
                         Т
                                           JP 2006-524091
                                                                  20040820
    NO 2006000719
                         Α
                               20060427
                                           NO 2006-719
                                                                  20060214
PRIORITY APPLN. INFO.:
                                           US 2003-496974P
                                                             P 20030821
                                                             A 20040819
                                           US 2004-923538
                                                             W 20040820
                                           WO 2004-US27195
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OTHER SOURCE(S): MARPAT 142:316496

GI

AB Title compds. I [Ring B = saturated or partially unsatd., (un) substituted cycloalkyl or heterocycle; X = O or S; Z = CO, CONR8, NR8, NR8CO, etc.; R1 = H, (un) substituted-alkyl, -alkenyl, -aryl, etc.; R2 = (un) substituted aryl or heteroaryl; R3 = H, Me, or Et; R8 = H, alkyl, or cycloalkyl; R10 and R10a independently = H or (un) substituted alkyl; R11 = H, alkyl, etc.; R12 = H, alkyl, (un) substituted carbocycle; m = O-1; n = 1 or 2], or pharmaceutically acceptable salt forms thereof, are prepared and disclosed as modulators of chemokine receptor activity. Thus, e.g., II was prepared by amidation of trans-4-aminocyclohexanol hydrochloride with (3-trifluoromethylbenzovlamino)acetic acid followed by meylation, substitution

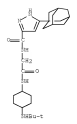
trifluoromethylbenzoylamino)acetic acid followed by mesylation, substitution with sodium azide and subsequent reduction I were deemed active (IC50 value of 20 µM or less) in antagonism of MCP-1 binding to human peripheral blood mononuclear cells. As modulators of MCP-1, I should prove useful for the prevention of asthma, multiple sclerosis, artherosclerosis, and rheumatoid arthritis.

IT 847953-11-1P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (USes)

(preparation of cycloalkylamine derivs, with chemokine receptor activity) RN $\,\,$ 847953-11-1 CAPLUS

CN 1H-Pyrazole-3-carboxamide, N-[2-[[4-[(1,1-dimethylethyl)amino]cyclohexyl]a mino]-2-oxoethyl]-5-tricyclo[3.3.1.13,7]dec-1-v1- (CA INDEX NAME)



L3 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:965056 CAPLUS Full-text

DOCUMENT NUMBER: 141:411216

TITLE: Preparation of amino acid pyrazolecarboxamides as heat shock protein 90 (HSP90) inhibitors for the treatment

of cancer
INVENTOR(S): Barril-Alonso, Xavier; Dymock, Brian William;

Drysdale, Martin James

PATENT ASSIGNEE(S): Vernalis Cambridge Limited, UK; Cancer Research
Technology Ltd.; The Institute of Cancer Research

SOURCE: PCT Int. Appl., 33 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE ---------WO 2004096212 A1 20041111 WO 2004-GB1740 20040423 WO 2004096212 A9 20050331 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN,

TD. TG EP 1620090 20060201 EP 2004-729149 20040423 A1 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, FI, RO, CY, TR, BG, CZ, EE, HU, PL, SK JP 2006524673 T 20061102 JP 2006-506168 20040423 US 2007072855 A1 20070329 US 2006-553955 20060810 PRIORITY APPLN. INFO.: GB 2003-9637 A 20030428

WO 2004-GB1740 W 20040423

Use of title compds. [I; R1 = Ar1(Alk1)pZr(Alk2)sQ; Ar1 = (substituted) arvl. AΒ heteroaryl; Alk1, Alk2 = alkylene, alkylene; p, r, s = 0, 1; Z = 0, S, CO, CS, SO2, CO2, CONRa, NRa, etc.; Ra = H, alkyl; Q = H, (substituted) carbocyclyl, heterocyclyl: R2 = Arl(Alkl)pZr(Alk2)sO, carboxamide, (substituted) carbocyclyl, heterocyclyl; R3 = H, (hydroxy-substituted) Me, Et, Pr; X = OR4, NR4R5; R4, R5 = H, (substituted) alkvl; NR4R5 = 5-8 membered heterocyclvl], for inhibition of HSP90 activity is claimed. Thus, 6-chloro-7-hydroxy-3-(4methoxyphenyl)-4-oxo-4H-chromene-2- carboxylic acid (preparation given) was heated with N2H4.H2O and aqueous NaHCO3 in EtOH at 70° for 2 h to give 5-(5chloro-2,4-dihydroxyphenyl)-4-(4- methoxyphenyl)-2H-pyrazole-3-carboxylic acid. This was stirred overnight with 1-hydroxybenzotriazole hydrate, Nethyl-N'-(3- dimethylaminopropyl)carbodiimide hydrochloride, Nmethylmorpholine, and racemic alanine Me ester hydrochloride in CH2Cl2 at 0° to room temperature to give racemic alanine Me ester 5-(5-chloro-2,4dihydroxyphenyl)-4- (4-methoxyphenyl)-2H-pyrazole-3-carboxamide. The latter in a fluorescence polarization assay showed IC50 <10 µM for binding to HSP90.

791103-68-9P 791103-69-0P 791103-70-3P 791103-71-4P 791103-72-5P 791103-73-6P 791103-74-7P 791103-76-9P 791103-77-0P

791103-78-1P 791103-79-2P 791103-80-5P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of amino acid pyrazolecarboxamides as heat shock protein 90 (HSP90) inhibitors for the treatment of cancer)

RN 791103-68-9 CAPLUS

CN Alanine, N-[[5-(5-chloro-2,4-dihydroxyphenyl)-4-(4-methoxyphenyl)-1Hpyrazol-3-yl]carbonyl]-, methyl ester (CA INDEX NAME)

RN 791103-69-0 CAPLUS

CN Alanine, N-[[5-(5-chloro-2,4-dihydroxyphenyl)-4-(4-methoxyphenyl)-1Hpyrazol-3-yl]carbonyl]- (CA INDEX NAME)

RN 791103-70-3 CAPLUS

CN Glycine, N-[[5-(5-chloro-2,4-dihydroxyphenyl)-4-(4-methoxyphenyl)-1Hpyrazol-3-yl]carbonyl]-, methyl ester (CA INDEX NAME)

RN 791103-71-4 CAPLUS

CN D-Alanine, N-[[5-(5-chloro-2,4-dihydroxyphenyl)-4-(4-methoxyphenyl)-1Hpyrazol-3-yl]carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 791103-72-5 CAPLUS

CN L-Alanine, N-[[5-(5-chloro-2,4-dihydroxyphenyl)-4-(4-methoxyphenyl)-1Hpyrazol-3-yl]carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

RN 791103-73-6 CAPLUS

CN 1H-Pyrazole-3-carboxamide, N-[(1R)-2-amino-1-methyl-2-oxoethyl]-5-(5-chloro-2,4-dihydroxyphenyl)-4-(4-methoxyphenyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 791103-74-7 CAPLUS

CN 1H-Pyrazole-3-carboxamide, N-[(1S)-2-amino-1-methyl-2-oxoethyl]-5-(5-chloro-2,4-dihydroxyphenyl)-4-(4-methoxyphenyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 791103-76-9 CAPLUS

CN 1H-Pyrazole-3-carboxamide, 5-(5-chloro-2,4-dihydroxyphenyl)-4-(4-methoxyphenyl)-N-((1R)-1-methyl-2-(methylamino)-2-oxoethyl]- (9CI) (CA INDEX NAME)

RN 791103-77-0 CAPLUS

CN 1H-Pyrazole-3-carboxamide, 5-(5-chloro-2,4-dihydroxyphenyl)-4-(4-methoxyphenyl)-N-(1R)-1-methyl-2-[(1-methylethyl)amino]-2-oxoethyl-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 791103-78-1 CAPLUS

CN 1H-Pyrazole-3-carboxamide, 5-(5-chloro-2,4-dihydroxyphenyl)-N-[(1R)-2-[[2-(dimethylamino)ethyl]amino]-1-methyl-2-oxoethyl]-4-(4-methoxyphenyl)-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 791103-79-2 CAPLUS

CN 1H-Pyrazole-3-carboxamide, 5-(5-chloro-2,4-dihydroxypheny1)-4-(4-methoxypheny1)-N-[(1R)-1-methy1-2-[[2-(4-morpholiny1)ethy1]amino]-2-oxoethy1]- (9CI) (CA INDEX NAME)

RN 791103-80-5 CAPLUS

CN 1H-Pyrazole-3-carboxamide, 5-(5-chloro-2,4-dihydroxyphenyl)-N-[(1R)-2-(ethylamino)-1-methyl-2-oxoethyl]-4-(4-methoxyphenyl)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 791103-83-8P 791103-84-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of amino acid pyrazolecarboxamides as heat shock protein 90 (HSP90) inhibitors for the treatment of cancer)

RN 791103-83-8 CAPLUS

CN D-Alanine, N-[[5-[5-chloro-2,4-bis[[(1,1-dimethylethoxy)carbonyl]oxy]pheny 1)-4-(4-methoxyphenyl)-1H-pyrazol-3-yl]carbonyl]-, methyl ester (9CI) (CA INDEX NAME)

791103-84-9 CAPLUS RN

CN D-Alanine, N-[[5-[5-chloro-2,4-bis[[(1,1-dimethylethoxy)carbonyl]oxy]pheny 1]-4-(4-methoxyphenyl)-1H-pyrazol-3-yl]carbonyl]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 4 OF 6 CAPLUS COPYRIGHT 2008 ACS on STN 2004:675727 CAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 141:207521

TITLE: Preparation of bis(hetero)aryl carboxamides as PGI2

antagonists for the treatment of urological disorders. INVENTOR(S): Murata, Toshiki; Shintani, Takuya; Umeda, Masaomi;

Lino, Takashi; Moriwaki, Toshiya

PATENT ASSIGNEE(S): Bayer Healthcare A.-G., Germany SOURCE: PCT Int. Appl., 101 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	TENT :				KIN		DATE									ATE	
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		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI
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SOURCE(S): 1:20752

GT

AB Title compds. I [X = -Ar1-Ar2-R1; Ar1, Ar2 = Ph, 5 or 6-membered heteroarom. ring containing 1-4 heteroatoms, e.g., O, N, S; R1 = OR11, SR11, SR11,

742057-80-3P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of bis(hetero)aryl carboxamides as PGI2 antagonists for the treatment of urol. disorders.)

RN 742057-80-3 CAPLUS

CN L-Phenylalanine, N-[[5-[4-(phenylmethoxy)phenyl]-1H-pyrazol-3-yl]carbonyl]-(CA INDEX NAME)

Absolute stereochemistry.

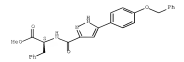
IT 742058-29-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of bis(hetero)aryl carboxamides as PGI2 antagonists for the treatment of urol. disorders.)

RN 742058-29-3 CAPLUS

CN L-Phenylalanine, N-[[5-[4-(phenylmethoxy)phenyl]-1H-pyrazol-3-yl]carbonyl], methyl ester (CA INDEX NAME)



L3 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER:

1986:406505 CAPLUS Full-text

DOCUMENT NUMBER: 105:6505 ORIGINAL REFERENCE NO.: 105:1213a,1216a

Phenylpyrazole derivatives

INVENTOR(S): Isekawa, Junichi; Shaku, Kunio; Sawada, Masahiro;

> Fukuda, Minoru; Yamada, Toshihiro; Oki, Masahiko; Matsuo, Yoshio

PATENT ASSIGNEE(S): Morishita Pharmaceutical Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF DOCUMENT TYPE: Patent

LANGUAGE: Japanese FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 61040266	A	19860226	JP 1984-163327	19840801
PRIORITY APPLN. INFO.:			JP 1984-163327	19840801
GI				

- AB The title compds. [I; R = H, lower alkyl, (un)esterified carboxyalkyl, Ph, PhCh2; R1 = (un)esterified CO2H, carbamoyl, (un)substituted lower alkyl; R2 = alkyl, (4- or α -substituted) PhCH2], useful as hypolipemics (no data), were prepared Thus, cyclocondensation of p- ClC6H4CH2OC6H4COCH:C(OMe)CO2Et-p with N2N4.H2O in refluxing EtOH for 1 h gave 70% I (R = H, R1 = CO2Et, R2 = 4-C1C6H4CH2).
- 102669-19-2P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as hypolipemic)
- 102669-19-2 CAPLUS RN CN Glycine, N-[[5-[4-[(4-chlorophenyl)methoxy]phenyl]-1H-pyrazol-3vl]carbonvl]-, ethvl ester (CA INDEX NAME)

L3 ANSWER 6 OF 6 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1970:415245 CAPLUS Full-text

DOCUMENT NUMBER: 73:15245

ORIGINAL REFERENCE NO.: 73:2553a,2556a

TITLE: Substances with antineoplastic activity. XLIII.

Reaction of ethyl ester of N-[β -(4-

methoxybenzoyl)- β -bromoacryloyl]glycine and $-\beta$ -alanine with hydrazide; ethyl ester of

 $N-[\beta-(4-methoxybenzoy1)-\beta-$

bromoacrylovljqlycylqlycine

AUTHOR(S): Zikan, Viktor; Semonsky, Miroslav; Svatek, Emil CORPORATE SOURCE: Vyzk. Ustav Farm. Biochem., Prague, Czech.

SOURCE: Vyzk. Ustav Farm. Biochem., Frague, Czech.

Collection of Czechoslovak Chemical Communications

(1970), 35(5), 1434-9

CODEN: CCCCAK; ISSN: 0010-0765

DOCUMENT TYPE: LANGUAGE:

Journal English

AB p-MeOC6H4COCBr:CHCONH(CH2)nCO2Et (n = 1 and 2) gave pyrazoles with excess N2H4.H2O in EtOH. p-MeOC6H4COCBr:CHCONHCH2CO2H gave with H2NCH2CO2Et by the dicyclohexylcarbodiimide method p-MeOC6H4COCBr:CHCONHCH2COCHECTOCHHCH2CO2Et (I), which exists predominantly in the hydroxylactam form. One of the pyrazoles and I inhibited the growth of the mammary adenocarcinoma, Ehrlich ascites tumor, and Crockers sarcoma 180 by 33-47% in rats but did not prolong survival of the animals. I prolonged the survival of mice with the \$ 37 sarcoma by 24% but had no effect on the tumor growth. None of the compds. had any effect on the Yoshida ascites sarcoma.

IT 27069-13-2P 27069-15-4P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 27069-13-2 CAPLUS

CN Glycine, N-[[5-(p-methoxyphenyl)pyrazol-3-y1]carbonyl]-, hydrazide (8CI)
 (CA INDEX NAME)

RN

CN Glycine, N-[[5-(p-methoxyphenyl)pyrazol-3-yl]carbonyl]- (8CI) (CA INDEX NAME)

=> =>

---Logging off of STN---

=>

Executing the logoff script...

=> LOG Y

COST IN U.S. DOLLARS	SINCE FILE	TOTAL
	ENTRY	SESSION
FULL ESTIMATED COST	58.14	236.92
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE	TOTAL
	ENTRY	SESSION
CA SUBSCRIBER PRICE	-4.80	-4.80

STN INTERNATIONAL LOGOFF AT 14:41:02 ON 11 FEB 2008